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(54) Magnetite particles and production process of the same

Magnetitteilchen und Verfahren zu ihrer Herstellung Particules de magnétite et leur procédé de préparation

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Description

[0001] The present invention relates to magnetite particles and a production process of the same, and more specifically to magnetite particles having an average particle size of not more than 0.6 μ m, a coercive force of 40 to 100 Oe (3.18 to 7.96 kA/m) at 10 kOe and electrical resistance of not less than 1 x 10⁴ Ω · cm in a particle shape of an octahedron and presenting a black color, which is particularly used as a use for an electrostatic copying magnetic toner magnetic powder, and a paint black pigment powder, and a production process of the same.

S. Description of the prior Art

[0002] Recently, as a magnetic toner material for a dry-type electronic copying machine, a printer or the like, magnetite particles which are obtained by the aqueous solution reaction process have widely been used. Various types of general development properties required for a magnetic toner. However, with the development of the electrophotographic properties has become higher. That is, a conventional character as well as the output of the photograph paper or photograph or the like are required. In particular, among the printer having an ability of 600 dots or more per inch has appeared so that a latent image on a photoreceptor becomes further precise. Accordingly, a high fine-line reproducibility in the development is strongly demanded. However, the Japanese Patent Application Laid-Open Gazette (Kokai) No. Hei 3-1 160, describes that toners are dispersed in the periphery of an image in a case where the coercive force of the magnetic powder exceeds 100 Oe, and that for this reason, it is difficult to respond to a demand for a higher teproducibility of a mesh point or tine line. Further, it seems that the powder having a higher electrical resistance is better in order to stabilize the charging of toner particles in a proper amount thereof. Such describes a toner which contains silicon and magnetic powders having electrical resistance of 1 x 10⁴ Ω - cm or more.

contains silicon and magnetic powders having electrical resistance of $1 \times 10^4 \,\Omega_{\odot}$ cm or more. [0004] In all of these above-mentioned publications, spherical magnetite particles are used. Magnetite particles, which satisfy a coercive force of 100 Oe or less, are approximately spherical while depending on the particle diameter. It is known that magnetite particles having a particle shape of, for example, an octahedron, have a coercive force of above 100 Oe, in an average particle diameter of not more than 0.6 μ m. The electrical resistance in magnetite particles

is usually 10² to 10³ Ω • cm.

SUMMARY OF THE INVENTION

[0005] Accordingly, it is an object of the present invention is to remove such conventional problems and provide magnetite particles with improved dispersibility, having an average particle diameter of a predetermined level or less, a coercive force of 100 Oe or less in a particle shape of an octahedron, and electrical resistance of not less than 1 x 10⁴ cive force of 100 Oe or less in a particle shape of an octahedron, and electrical resistance of not less than 1 x 10⁴

 $\Omega \cdot cm$, and further provide a production process of the magnetite particles. The present inventors have tried to strain the lattice of the magnetite crystal by allowing silicon to contain inside

octahedron-shaped magnetite particles.

[0007] The B-H hysteresis loop of the silicon contained magnetite particles therein is characteristically bent as shown in Fig. 1. It seemed that the bending of this B-H hysteresis loop is derived from a magnetic anisotropy based on the crystal lattice strain. Thus, we have found that when relaxation of the lattice strain by heat treatment is evaluated, the crystal lattice strain. Thus, we have found that when relaxation of the lattice strain by heat treatment is evaluated, the

partide coercive force is decreased.

[0008] Each of the magnetite particles according to the present invention has an octahedron shape. The octahedron-shaped ones, shaped magnetite particles are advantageous in that they have larger stable regions than the spherical-shaped ones, so that the former is more easily produced than the latter in the production process, and that the distribution of the particle so that the former is more easily produced than the latter in the production process, and that the distribution of the particle so that the former is more easily produced than the latter in the production process, and that the distribution of the particle so that the former is more easily produced than the latter in the production process, and that the distribution of the particle so that the former is more easily produced than the latter in the production process, and that the distribution of the particle so that the former is more easily produced than the latter in the production process.

ticle size of the octahedron-shaped magnetite particle is sharp.

The magnetite particles according to the present invention are characterized in that the particle shape is octahedron, the average particle diameter is not more than 0.6 µm, the coercive force is 40 to 100 Oe (3.18 to 7.96 kA/m)

at 10 kOe, and the electrical resistance is not less than 1 x 10 4 Ω \cdot cm. [0010] While the novel features of the present invention are set forth particularly in the appended claims, the invention will be better understood and appreciated, along with other objects and features thereof, from the following detailed

description taken in conjunction with the attached drawing.

BRIEF DESCRIPTION OF THE DRAWING

[0011] Fig. 1 is a view showing a hysteresis loop (the external magnetic fields of 1 kOe and 10 kOe) of magnetite particles described in Example 1.

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PREFERRED EMBODIMENTS OF THE INVENTION

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[0012] The average particle diameter of the magnetite particles according to the present invention is not more than 0.6 μ m, preferably 0.1 to 0.6 μ m. When the average particle diameter is too small, the magnetite particles tend to agglomerate and the environment resistance of the particles is deteriorated. Further, the degree of black color is also deteriorated as the hue thereof. Furthermore, in such case, the magnetite particles has superparamagnetism, whereby the coercive force as well as the saturation magnetism are decreased. In a case where the average particle diameter is too large, when the magnetite particles dispersed into the toner are used, it is excessively stuck out of the surface of the toner or the deviation of the particle occurs, which is undesirable. In other words, since the magnetite particles have a single magnetic domain structure in the average particle diameter of about not more than 0.18 μ m, the coercive force of the magnetite particle becomes large. However, when the magnetite particles have a smaller average particle diameter than the above-mentioned one, magnetite particles which present the superparamagnetism is increased (DENKI KAGAKU (Japan), Vol. 37, p 330 by Takada).

[0013] When the coercive force exceeds 100 Oe, toners are dispersed in the periphery of an image as mentionedabove, and for this reason, it is difficult to respond to a demand for a higher reproducibility of a mesh point or fine line. Further, it is substantially difficult to produce magnetite particles having a coercive force of less than 40 Oe.

[0014] Further, if the electrical resistance of the magnetite particles is 10³ order or less, it is often difficult to maintain the charging stability in the toners.

[0015] Further, the magnetite particles according to the present invention is characterized in that they contain silicon of 0.1 to 3.0% by weight, with respect to the magnetite particles, and a total 0.5 to 4.5% by weight of components of Si, Al, Ti, Zr, Mn, Mg and Zn other than Fe in the time when the Fe is dissolved by 20% from the surface of the particles, with respect to the magnetite particles.

[0016] If the silicon contents are less than 0.1% by weight, a coercive force within the above-mentioned range cannot be obtained. On the other hand, if the silicon contents are over 3.0% by weight, much silicon is often exposed on the surface of the magnetite particles, whereby the working properties become bad.

[0017] Further, if the total contents of components of Si, Al, Ti, Zr, Mn, Mg and Zn other than Fe in the time when the Fe is dissolved by 20% from the surface of the particles, are less than 0.5% by weight with respect to the magnetite particles, effects on the increase in the electrical resistance are small. On the other hand, the total contents thereof are over 4.5% by weight, the saturation magnetism is undesirably decreased.

[0018] Next, a production process of the magnetite particles according to the present invention will be described.

[0019] The production process of the magnetite particles according to the present invention comprises the steps of mixing a ferrous salt aqueous solution with a silicate aqueous solution to obtain an aqueous mixture solution, oxidizing at not less than pH 10 after mixing the aqueous mixture solution with an alkaline aqueous solution to obtain a slurry, adding an aqueous solution containing at least one component selected from the group consisting of Si, Al, Ti, Zr, Mn, Mg, Zn and Fe into the slurry when the oxidation reaction is progressed by 80% or more, to prepare the mixture so that it exhibits pH 5 or more, further oxidizing the mixture to obtain an oxidized slurry containing magnetite particles, and heat treating the magnetite particles after filtrating, washing, drying and grinding the thus obtained slurry, so that the contents of FeO within the magnetite particles obtained by oxidation is 18% by weight or more.

[0020] According to the present invention, a ferrous salt aqueous solution is mixed with a silicate aqueous solution. As the ferrous salt aqueous solution, a ferrous sulfate aqueous solution or the like may be given. The concentration and amount of the silicate aqueous solution are prepared so that silicon of 0.1 to 3.0% by weight is contained with respect to a finally obtained magnetite particles.

[0021] After that, the mixed solution of this ferrous salt aqueous solution and the silicate aqueous solution is mixed with an alkaline aqueous solution to form iron hydroxide slurry and an oxygen containing gas, preferably the air is blown into to carry out the oxidation reaction at a temperature of 60 to 100°C, preferably, 80 to 90°C.

[0022] An aqueous solution which contains at least one of components consisting of Si, Al, Ti, Zr, Mn, Mg, Zn and Fe is poured into slurry when the oxidation reaction is progressed by 80% or more, that is, by 80 to 100%, and the oxidation is repeated.

[0023] As the selected components in this process, even an acid type, such as sulfuric acid, or an alkaline type can be used. Further, even a simultaneous addition of the selected components or separated addition thereof to the slurry may be carried out.

[0024] After the oxidation reaction is completed, each of processes of washing, filtration, drying and grinding is carried out to obtain magnetite particles. This drying process means that the moisture within the cake is evaporated at a temperature of below 100°C and then, a loss of the moisture within the powder reaches 0.5% or less at a temperature of 100°C for an hour.

[0025] After that, according to the present invention, this magnetite particles are heat treated. As the atmosphere of the heat treatment, any of an oxidizing atmosphere such as the air or the like and an inert atmosphere such as the nitrogen or the like can be used. Further, although various heat treatment temperatures are used in accordance with the

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[0026] It is necessary to prevent the deterioration of the degree of a black color in the heat treatment of the magnetite son with the coercive force of a dried, non-heated magnetite particle by this heat treatment. ticle for a suitable time. The coercive force of the magnetite particles is further decreased by 10% or more in comparimay be selected so that the heat treatment can be carried out at below the sintering temperature of the magnetite par-

to 300°C. On the other hand, when the heat treatment is carried out in an inert atmosphere, a heat treatment process ment is industrially difficult. Thus, it is preferable that the heat treatment temperature of the magnetite particles is 100 treatment is carried out at a high temperature, the heat treatment time becomes short. However, the short time treatment is carried out at a low temperature, the heat treatment time becomes long, and on the other hand, when the heat may be set so that FeO within the magnetite particles is not less than 18% by weight. Nevertheless, when the heat treatmagnetite particle size or time, when the heat treatment is carried out in an oxidizing atmosphere, the heat temperature

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that the coercive force thereof is increased in the approximate composition of FeO = 10%. This phenomenon is are then oxidized, so that the color of the magnetite particle is gradually changed from black to brown with the result ventionally been recognized that when the magnetite particles are gradually oxidized Fe^{24} ions within Fe_3O_4 particles partide by setting the temperature and the time so that FeO within the magnetite particle is 18% or more. It has con-

[0027] The present inventors have assumed that since the coercive force of the magnetite particles was decreased explained by the fact that a part of Fe₃O₄ particles is changed to a γ - Fe₂O₃ intermediate oxide film.

[0028] Furthermore, when the heat treatment is carried out, the saturation magnetization (5s) tended to increase par-50 be slightly curved. from heat relaxation of the lattice strain. Further, the B-H hysteresis loop of heat treated magnetite particles tended to regardless of an oxidizing or an inert heat treatment atmosphere, the decrease in the coercive force thereof is derived

based on the lattice strain is improved to thereby enhance the magnetic permeability by the heat treatment. ticularly in the external magnetic field of 1 kOe. It seems that this is because the magnetic anisotropy which is mainly

ing the selected components on the surface of the powders in the presence of non-reacted Fe rather than treating only present inventors have found that the increase in electrical resistance is effected by forming an iron oxide layer containsisting of Si, Ai, Ti, Zr, Mn, Mg and Zn to contain into the surface layer thereof cannot be understood. However, the [0029] The reasons why an electrical resistance of powders is increased by allowing at least one of components con-

the selected components thereon. 52

present invention have such features that they are well dispersed into toners, with the result that they are preferably with the selected components, and performing a heat treatment. Accordingly, the magnetite particles according to the less than 1 x 10⁴ Ω \cdot cm, by allowing silicon to contain into the magnetite particles, improving the surface of the particles shape of an octahedron, a coercive force of 40 to 100 Oe (3.18 to 7.96 kA/m) at 10 kOe, and electrical resistance of not it is possible to obtain magnetite particles having an average particle diameter of not more than 0.6 µm in a particle [0030] As described previously and apparent from the following disclosure, in accordance with the present invention,

used for particularly, electrostatic copying magnetic toners.

erence to the attached drawing. [10031] In the following paragraphs, the present invention will be described in more detail by way of examples with ref-

Example 1

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(1) Particle diameter

subjected to a usual washing, filtration, drying and grinding to thereby obtain magnetite particles. prepared so that it exhibits pH 5 or more. After that, reoxidation was completely carried out. The obtained particles were of the respective aqueous solutions containing 0.84 mol/l And 2.09 mol/l Fe were added to the obtained mixture, and blown into the mixture while keeping the temperature at 90°C, and the reaction was allowed to complete. After that, 3 I 50 l of 3.58 mol/s sodium hydroxide while stirring it, so that the mixture exhibits pH 10 or more, and 20 l/min of air was an aqueous solution containing 1.62 mol/l Si to obtain a mixture. After that, the thus obtained mixture was mixed with [0032] As shown in Table 1, 48 I of ferrous sulfate aqueous solution containing 1.87 mol/l Fe^{2+} was mixed with 2 I of

shown in Table 2. The magnetic hysteresis loop (1 kOe, and 10 kOe of external magnetic fields) of the magnetite partities, the electrical resistance and the dispersibility were evaluated by the following methods. The obtained results are [0033] As to the obtained magnetite particles, the particle diameter, the specific surface area, the magnetic proper-

cles is shown in Fig. 1.

(30000 x magnification) of the particles and the average thereof was defined as a particle diameter [0034] Some of particle diameters on a picture were measured by using a transmission electron microscopic picture

(2) Specific surface area

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[0035] 2200 type BET meter produced by Shimazu-Micromerytics was used.

(3) Magnetic properties

[0036] VSM-P7 type of vibration specimen type magnetometer produced by Touei kougyo was used.

(4) Electrical resistance

[0037] After 10 g of a sample is placed in a holder, a pressure of 600 Kg/cm² is applied to the holder to mold into a 25 mm ϕ tablet sample. After that, an electrode is set on the sample, measurement is carried out while pressing it at a pressure of 150 Kg/cm². The respective values of electrical resistance of magnetite particles were obtained by calculating from the thickness, cross-sectional area and value of electrical resistance of the samples.

(5) Dispersibility

[0038] After 1 g of magnetite and 0.7 g of linseed oil were mixed and kneaded, 4.5 g of clear lacquer was added to the mixture and further kneaded it. This mixture was applied on a glass plate by using an 4 mil applicator. After drying the applied mixture, reflectance at 60° were measured by the Murakami type GLOSS METER (GM-3M).

(6) Analysis of surface layer

[0039] After 5 I of 1 N hydrochloric acid was poured into a 25 g sample to obtain a mixture, the mixture was stirred and dissolved at 50°C. The thus dissolved mixture solution was filtered by using a membrane filter having a pore size of 0.1 µm to obtain a filtrate. The thus obtained filtrate was analyzed and the analysis values of the components other than Fe at the time when 20% Fe was dissolved were calculated by percentage by weight in the magnetite respectively.

Examples 2 to 7

[0040] As shown in Table 1, items other than the amount of added Si which is not reacted with other compositions, the addition time for the surface layer, the types of additives and the addition amount were used in the same manner as in Example 1, and magnetite particles were obtained. The obtained magnetite particles were evaluated by the same methods as in Example 1. The obtained results are shown in Table 2.

Comparative Example 1

[0041] As shown in Table 1, the same manner as in Example 1 was carried out without adding silicon and surface components before the reaction, and magnetite particles were obtained. The obtained results were evaluated in the same manner as in Example 1. The results are shown in Table 2.

Comparative Example 2

[0042] As shown in Table 1, the same manner as in Example 3 was carried out except for without adding silicon before the reaction, and magnetite particles were obtained. The obtained results were evaluated in the same manner as in Example 1. The results are shown in Table 2.

Comparative Example 3

[0043] As shown in Table 1, the same manner as in Example 5 was carried out except for without adding surface components before the reaction, and magnetite particles were obtained. The obtained results were evaluated in the same manner as in Example 1. The results are shown in Table 2.

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|-------------|-------------|-------------|-------|-------|-------|-------|-------|-------|-------|---------|-----------------------|--------------------------------------|
| Comp. Ex. 3 | Comp. Ex. 2 | Comp. Ex. 1 | Ex. 7 | Ex. 6 | Ex. 5 | Ex. 4 | Ex. 3 | Ex. 2 | Ex. 1 | | | |
| 1.87 | 1.87 | 1. 87 | 1. 87 | 1. 87 | 1. 87 | 1. 87 | 1. 87 | 1. 87 | 1. 87 | (mol/1) | Fe concentration | Ferrous sulfate aqueous solution |
| 48 | 48 | 48 | 48 | 48 | 48 | 48 | 48 | 48 | 48 | (1) | liquid amount | fate ution |
| 2. 03 | - | | 3. 38 | 2. 71 | 2. 03 | 2. 03 | 1. 62 | 1. 09 | 1. 62 | (mol/1) | Si concentration | Silicon containing aqueous solution |
| 2 | 1 | į | 2 | 2 | 2 | 2 | 2 | 2 | 2 | (1) | liquid amount | aining tion |
| 3. 58 | 3. 58 | 3. 58 | 3. 58 | 3. 58 | 3. 58 | 3. 58 | 3.58 | 3.58 | 3.58 | (mol/1) | NaON concentration | Sodium hydroxide aqueous solution |
| 50 | 50 | 50 | 20 | 50 | 50 | 50 | 50 | 50 | 50 | (E) | liquid amount | xide tion |
| 11.2 | 11.4 | 11.5 | 11. 3 | 10.8 | 11.1 | 11.7 | 11. 4 | 11. 3 | 11.5 | | 1 | B |
| | 90 | 1 | 83 | 88 | 93 | 95 | 90 | 94 | 100 | | ratio) (%) | Addition time (Reation |

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(continued)

| | Conc | entration | Concentration of components in the aqueous solution (3L) added | nts in the | adneons s | olution (3 | L) added | |
|-------------|---------|-----------|--|------------|-----------|------------|----------|---------|
| | Si | ΙV | Ti | JZ | уļu | Mg | Zn | Fe |
| | (mol/l) | (mol/l) | (mol/l) | (mol/l) | (mol/l) | (mol/1) | (mol/1) | (mol/1) |
| Ex. 1 | 1 | 0.84 | | 1 | | _ | - | 2.09 |
| Ex. 2 | | 0.93 | 0.26 | l | | _ | 0.19 | 0.91 |
| 5x. 3 | 0.45 | 0.47 | ı | 1 | | ï | | 1 |
| Bx. 4 | l | - | ı | l | 0.69 | _ | 1 | ı |
| Ех. 5 | 0.45 | ١ | 1 | _ | 0. 46 | l | _ | |
| Ex. 6 | 0.27 | 0.75 | l | l | 1 | 0.32 | | |
| Ex. 7 | | 0.94 | 1 | 0.14 | 0.92 | 1 | 1 | 1 |
| Comp. Ex. 1 | | I | ١ | ı | | l | | 1 |
| Comp. Ex. 2 | 0. 45 | 0.47 | | _ | 1 | l | 1 | |
| Comp. Ex. 3 | 1 | 1 | ١ | - | 1 | 1 | 1 | 1 |

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[0044] It has been appreciated from Examples 1 to 7 and Comparative Examples 1 to 3 that magnetite particles having an average particle diameter of not more than 0.6 µm has a coercive force of not less than 1 00 Oe.

[0045] As can be seen from Comparative Examples 1 and 3, the magnetite particles which have not added metals on

| į | | (10kue) | | | (1k0e) | | e | tance | 8 | 8 | values (%) |
|-----------------------------------|-------|---|---------------------------|---------------------------------------|--|---------------------------|---------|-------|-------|-----|------------------------------------|
| (mag) Satural magnet zation (cm// | tion | Residual magneti- zation (em \(\mu / \text{g} \) | Coercive force (0e) | Saturation magneti- zation (cm \mu/g) | Residual magneti- zation (em \(\mu / \(g \)) | Coercive force (Oc) | | | | | of components in the surface layer |
| Ex. 1 0.41 11 7 | 79. 7 | 7. 4 | 109 | 55. 9 | 6.2 | 92 | 1 x 10' | 90 | 27.0 | 1.2 | 1. 3 |
| Ex. 2 0.42 14 · 8 | 82.0 | 8. 2 | 107 | 57.0 | 7. 3 | 104 | 4 x 10° | 89 | 25. 9 | 0.8 | 2. 3 |
| Ex. 3 0.43 14 8 | 80. 9 | 6. 8 | 106 | 55. 9 | 6. 1 | 92 | 1 x 10° | 91 | 27. 5 | 1.7 | 1.4 |
| Ex. 4 0.52 12 8 | 81.3 | 7. 1 | 102 | 56. 9 | 6. 5 | 9.4 | 2 x 10* | 89 | 27. 8 | 1.5 | 1.9 |
| Ex. 5 0.25 16 8 | 82. 4 | 9. 1 | 117 | 58.7 | 8.0 | 108 | 1 x 10* | 89 | 26. 9 | 2.0 | 1. 9 |
| Ex. 6 0.19 18 8 | 80.7 | 9. 1 | 121 | 59.8 | 8. 6 | 110 | 2 x 10* | 90 | 26. 5 | 2.3 | 2. 0 |
| Ex. 7 0.39 21 7 | 79.3 | 7. 2 | 101 | 57. 2 | 6. 3 | 9. 8 | 6 x 10* | 89 | 26. 8 | 2.5 | 4.1 |
| Comp. 0.40 6 8 | 85. 4 | 10.5 | 117 | 62. 1 | 9. 8 | 108 | 8 x 10° | 80 | 26. 3 | 0 | 0. 3 |
| Comp. 0.41 13 8 | 82. 5 | 10.3 | 115 | 58. 3 | 9. 5 | 105 | 2 x 10* | 83 | 26. 8 | 0.5 | 1. 2 |
| Comp. 0.28 8 8 | 85. 0 | 9. 4 | 126 | 60.3 | 8. 6 | 117 | 4 x 10° | 84 | 27. 2 | 1.5 | 0.3 |

Table 2

a surface layer have the electrical resistance of less than $10^4 \,\Omega$ • cm. For Comparative Example 2, the value of the electrical resistance is $10^4 \,\Omega$ • cm order.

[0046] As can be seen from the results of reflectance, the magnetic particles that contain silicon therein and has a surface layer, have an improved dispersibility.

[0047] The respective particles were heat-treated by the following conditions.

Examples 8 to 10

[0048] Samples of Examples 1, 3 and 5 were heated at 150 °C in the air to prepare FeO. As to the obtained magnetite particles, the magnetic property, electrical resistance and reflectance were evaluated. The evaluated results are shown in Table 3.

Example 11

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[0049] A sample of Example 1 was heated at 300 °C for 30 min. in a nitrogen atmosphere. As to the obtained magnetite particles, the magnetic property, electrical resistance and reflectance were evaluated. The evaluated results are shown in Table 3.

Comparative Examples 4 to 6

[0050] Samples of Comparative Examples 1 to 3 were heated at 150 °C in the air to prepare FeO. As to the obtained magnetite particles, the magnetic property, electrical resistance and reflectance were evaluated. The evaluated results are shown in Table 3.

[0052] As can be seen from the results of Comparative Example 4, for a sample which does not contain silicon therein,

[0051] As can be seen from the results of Examples 8 to 11, the coercive force of the particles is reduced to be not a more than 100 Oe by heat treatment. Although the electrical resistance is slightly reduced in the inert gas atmosphere.

| | | | | | | | 3 | _ | D. FI |
|-------------|---|---------------------------------------|---------------------------|---------------------------------------|--|---------------------------|------------|-----------------------|-------------|
| | Mag | Magnetic properties (10k0e) | ties | Mag | Magnetic properties (1kOe) | ties | 3 3 | Electrical resistance | Keflectance |
| | Saturation magneti-zation (em \(\mu / g \) | Residual magneti-zation (em μ/g) | Coercive force (0e) | Saturation magneti- zation (cm \mu/g) | Residual magneti- zation (em \(\mu / g \) | Coercive force (0e) | | | |
| Ex. 8 | 79.4 | 6. 1 | 89 | 58. 6 | 6.2 | 89 | 23. 0 | 9 x 10° | 91 |
| Ex. 9 | 80.3 | 5. 0 | 76 | 58. 5 | 4. 5 | 70 | 26.0 | 3 x 10° | 91 |
| Ex. 10 | 80.1 | 6.9 | 80 | 62. 4 | 6.5 | 75 | 19.7 | 9 x 104 | 90 |
| Ex. 11 | 80. 6 | 5.7 | 83 | 60. 2 | 5. 2 | 78 | 26. 9 | 2 x 10° | 91 |
| Comp. Ex. 4 | 79.8 | 11. 2 | 125 | 60. 1 | 10.4 | 116 | 21.0 | 2 x 10° | 80 |
| Comp. Ex. 5 | 80.3 | 10.5 | 121 | 56. 9 | 10.1 | 116 | 21. 5 | 4 x 10* | 84 |
| Сопр. Ех. б | 81. 2 | 7. 3 | 85 | 61. 2 | 7. 4 | 83 | 20. 4 | 6 x 10* | 8.4 |

Table 3

the coercive force was increased by heat treatment.

it does not change in the air so as to be 10. Ω - cm order or more.

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[0053] As can also be seen from the results of Comparative Example 5, although the sample thereof has an electrical resistance of $10^4 \,\Omega$ cm order, the coercive force was increased by heat treatment, as in Comparative Example 4, due to an improved surface layer.

[0054] Further, as can be seen from the results of Comparative Example 6, since the sample contains silicon therein, the coercive force is reduced by heat treatment. However, since the surface layer is not improved, the electrical resistance is low.

[0055] As can be seen from these results, coercive forces of all magnetite particles containing silicon therein, in which lattices of the magnetite particles are allowed to strain were decreased. On the other hand, coercive forces of most silicon-free magnetite particles were slightly increased. Further, by improving the surface layer, the electrical resistance was enhanced and the dispersibility had an improved tendency.

[0056] Further, for magnetite particles containing silicon therein, a saturation magnetization tended to increase at 1 KOe in the external magnetic field, by heat treatment.

[0057] The present invention may be embodied in other specific forms without departing from the spirit or essential characteristics thereof. The present embodiment is therefore to be considered in all respects as illustrative and not restrictive, the scope of the invention being indicated by the appended claims rather than by the foregoing description and all changes which come within meaning and range of equivalency of the claims are therefore intended to be embraced therein.

Claims

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- Magnetite particles characterized in that the particle shape is octahedron, the average particle diameter is not more than 0.6 μm, the coercive force is 40 to 100 Oe at 10 kOe, and the electrical resistance is not less than 1 x 10⁴ Ω • cm.
- 25 2. Magnetite particles according to claim 1, wherein they contain silicon of 0.1 to 3.0% by weight, with respect to the magnetite particles, and a total 0.5 to 4.5% by weight of components of Si, Al, Ti, Zr, Mn, Mg and Zn other than Fe in the time when the Fe is dissolved by 20% from the surface of the particles, with respect to the magnetite particles.
- 30 3. A process for producing magnetite particles according to claims 1 and 2 comprising the steps of

mixing a ferrous salt aqueous solution with a silicate aqueous solution to obtain an aqueous mixture solution, oxidizing at not less than pH 10 after mixing the aqueous mixture solution with an alkaline aqueous solution to obtain a slurry,

adding an aqueous solution containing at least one component selected from the group consisting of Si, Al, Ti, Zr, Mn, Mg, Zn and Fe into the slurry when the oxidation reaction is progressed by 80% or more, to prepare the mixture so that it exhibits pH 5 or more,

further oxidizing the mixture to obtain an oxidized slurry containing magnetite particles, and

heat treating magnetite particles after filtrating, washing, drying and grinding the thus obtained slurry, so that the contents of FeO within the magnetite particles obtained by oxidation is 18% by weight or more.

4. A process for producing magnetite particles according to claim 3, wherein said heat treatment atmosphere is an oxidizing atmosphere or an inert atmosphere.

45 Patentansprüche

- Magnetitpartikel, dadurch gekennzeichnet, daß die Partikelform die eines Oktaeders ist, daß der mittlere Partikeldurchmesser nicht größer als 0,6 μm ist, daß die Koerzitivkraft 40 bis 100 Oe bei 10 KOe beträgt und daß der elektrische Widerstand nicht geringer als 1 x 10⁴ Ω • cm ist.
- 2. Magnetitpartikel nach Anspruch 1, worin in Bezug auf die Magnetitpartikel Silizium mit einem Gewichtsanteil von 0,1 bis 3,0% enthalten ist, und zum Zeitpunkt, zu dem 20% des Fe von der Oberfläche der Partikel aufgelöst ist, in Bezug auf die Magnetitpartikel, andere Bestandteile als Fe, wie Si, Al, Ti, Zr, Mn, Mg und Zn als Fe mit einem Gesamtgewichtsanteil von 0,5 bis 4,5% enthalten sind.
- 3. Ein Verfahren zum Herstellen von Magnetitpartikeln nach den Ansprüchen 1 und 2, das folgende Schritte umfaßt:

Mischen einer wässrigen Eisen(II)-Salzlösung mit einer wässrigen Silikatlösung, um ein wässriges Lösungs-

Eb 0 85e e32 B1

18% Gewichtsanteil oder mehr beträgt.

18% en poids ou plus.

Ben erhaltenen Breis, so daß der Gehalt an FeO innerhalb der über die Oxidation erhaltenen Magnetitpartikel Hitzebehandeln der Magnetitpartikel nach dem Filtrieren, Waschen, Trocknen und Zerkleinern des solchermaweiteres Oxidieren des Gemischs, um einen oxidierten Brei zu erhalten, der Magnetitpartikel enthält; und mehr fortgeschritten ist, um das Gemisch so verarbeiten, daß es einen pH-Wert von 5 oder mehr aufweist; ausgewählt ist, die aus Si, Al, Ti, Zr, Mn, Mg, Zn und Fe besteht, sobald die Oxidationsreaktion um 80% oder Hinzulügen einer wässrigen Lösung in den Brei, die mindestens einen Bestandteil enthält, der aus der Gruppe einer wässrigen alkalischen Lösung vermischt wurde, um einen Brei zu erhalten; Oxidieren bei einem pH-Wert der nicht weniger als 10 beträgt, nachdem das wässrige Lösungsgemisch mit gemisch zu erhalten;

oxidierende Atmosphäre oder eine inerte Atmosphäre ist. Ein Verlahren zum Herstellen der Magnetitpartikel nach Anspruch 3, worin die Hitzebehandlungsatmosphäre eine

Revendications 91

- 104 scm. 02 rieur à 0,6 µm, la force coercitive est de 40 à 100 Oe à 10 kOe et la résistance électrique n'est pas inférieure à 1 x caractérisées en ce que la forme des particules est octaédrique, le diamètre moyen des particules n'est pas supé-Particules de magnétite,
- sout de 20% à partir de la surface des particules par rapport aux particules de magnétite. 52 un total de 0,5 à 4,5% en poids de composants Si, Al, Ti, Zr, Mn, Mg et Zn autre que Fe au moment où Fe est disdans lesquelles elles contiennent du silicium de 0,1 à 3,0% en poids, par rapport aux particules de magnétite, et Particules de magnétite selon la revendication 1,
- Procédé pour produire des particules de magnétite selon les revendications 1 et 2, comprenant les étapes de :
- oxyder à pas moins de pH 10 après mélange, la solution aqueuse du mélange avec une solution aqueuse alcaadnense qn melange, mélanger une solution aqueuse de sel ferreux avec une solution aqueuse de silicate pour obtenir une solution
- Ti, Zr, Mn, Mg, Zn et Fe dans la bouillie quand la réaction d'oxydation a progressé de 80% ou plus, pour préajouter une solution aqueuse contenant au moins un composant choisi à partir du groupe comprenant Si, Al, line pour obtenir une bouillie,
- ainsi obtenue, de sorte que la teneur en FeO dans les particules de magnétite obtenues par oxydation est de traiter thermiquement les particules de magnétite après filtration, lavage, séchage et broyage de la bouillie oxyder encore le mélange pour obtenir une bouillie oxydée contenant des particules de magnétite, et parer le mélange de sorte qu'il montre le pH 5 ou plus,
- tement thermique est une atmosphère oxydante ou une atmosphère inerte. Procédé pour produire des particules de magnétite selon la revendication 3, dans lequel ladite atmosphère de trai-

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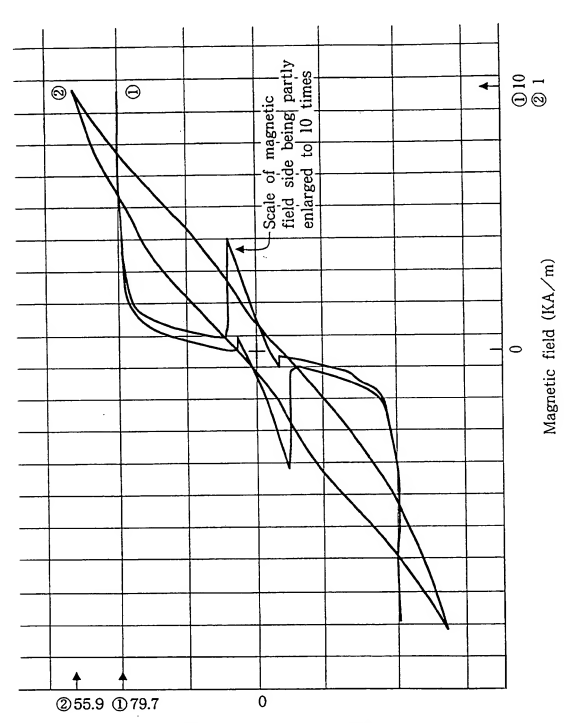
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Magnetization (Am²/Kg)

Fig. 1

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